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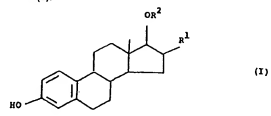
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(54) ESTRADIOL DERIVATIVES

(71) We, TAKEDA YAKUHIN KOGYO KABUSHIKI KAISHA, also known as TAKEDA CHEMICAL INDUSTRIES LTD., of 27 Doshomachi 2-chome, Higashi-ku, Osaka, Japan, a body corporate organised under the laws of Japan, do hereby declare the invention, for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following statement:—

The present invention relates to novel and useful 16β -alkylestradiol derivatives and to a process for producing them.

More particularly, the present invention relates to 16β -alkylestradiols represented by the formula (I):



wherein R^1 is an alkyl group or an alkenyl group of two or more carbon atoms; and R^2 is hydrogen or an acyl group (as herein defined), and to a process for producing the compounds (I).

Hitherto, testosterone or derivatives thereof (e.g. testosterone propionate) have been introduced for the therapy of estrogen-dependent disease (e.g. advanced breast cancer) as antiestrogen drugs. However, the therapy is generally accompanied with the drawback *inter alia* that the virilizing effect resulting from the androgenic potency of testosterone prevents the patient from continuing with the therapy.

We have discovered that 16β -alkylestradiol derivatives have substantially no estrogen activity but rather have an antiestrogen activity, and that this propensity is particularly pronounced where the number of carbon atoms in the 16β -alkyl moiety is within the range of from 2 to 4. The present invention is accomplished on the basis of these findings.

The present invention provides compounds of the general formula (I), which are useful as an antiestrogen drug, and a process for producing the compounds (I). Referring to the formula (I) and to formula (II) described below, the alkyl

Referring to the formula (I) and to formula (II) described below, the alkyl group or alkenyl group of two or more carbon atoms designated by R¹ may be straight-chain or branched, and saturated or unsaturated, thus being exemplified by lower alkyl groups having 2 to 4 carbon atoms, such as ethyl, propyl, isopropyl, butyl, isobutyl, sec-butyl, tert-butyl, allyl and 3-butenyl. The acyl group designated by R² in formula (I) above and by R² and R³ in formula (II) below is defined as a hydrocarbon-carbonyl group whose hydrocarbon moiety has from 1 to 8 carbon atoms. The hydrocarbon-carbonyl group is exemplified by lower alkylcarbonyl groups whose alkyl moieties have 1 to 3 carbon atoms, e.g. acetyl, propionyl, butyryl; arylcarbonyl groups, e.g. benzoyl; and aralkylcarbonyl groups, e.g. phenylpropionyl. Where R² and R² are an acyl group, the substituent —OR² or —OR² in the 17-position of formula (I) or (II) is an esterified hydroxyl group, and the corresponding compound is a 17-ester of the compound (I) or (II). The

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hydrocarbon radical designated by R² in formula (II) is an alkyl, aryl or aralkyl group. The alkyl group mentioned for R3 may be a straight-chain or branched lower alkyl group of 1 to 3 carbon atoms, viz. methyl, ethyl, propyl or isopropyl; the aryl group mentioned for R3 may, for example, be phenyl or p-nitrophenyl; and the aralkyl group for R² may, for example, be benzyl or benzhydryl.

The compounds (I) of the present invention can be produced according to per se known methods. For example, the compounds (I) may be produced according to the method illustrated as follows:

10 wherin R1 and R2 have the same meaning as defined above, R2' is hydrogen or an acyl group, and R3 is a hydrocarbon radical or an acyl group.

Thus, the above method is carried out by subjecting the compound (II) to a reaction leading to the cleavage of the acyl group or hydrocarbon radical of the esterified or etherified hydroxyl group in the 3-position thereof.

By the present reaction, the acyl group or hydrocarbon radical of the esterified or etherified hydroxyl group in the 3-position is removed, thus leaving a free

hydroxyl group in the 3-position. This reaction, where R3 is an alkyl or aryl group, that is to say where —OR3 is

an etherified hydroxyl group, is carried out by reacting the compound (II) with a reagent capable of cleaving an ether linkage. The ether-cleaving reagent may be any reagent which is able to cleave the ether linkage of the etherified hydroxyl group in the 3-position without affecting the steroid skeleton and the 16\beta-alkyl group of the starting compound. Thus, for example, there may be mentioned acidic reagents, for example, hydrohalic acids such as hydrochloric acid, hydrobromic acid and hydroiodic acid, halides of phosphorus, boron, aluminium, thallium and titanium, preferably the corresponding chlorides and bromides (e.g. phosphorus tribromide, boron tribromide, aluminium chloride, titanium tetrachloride), pyridinium halides (e.g. pyridinium chloride); Grignard reagents (e.g. methylmagnesium iodide and ethylmagnesium bromide); and sodium iodide dimethylsulfoxide. Generally, such ether-cleaving reagents are used in amounts within the range of from 1 to 10 moles per mole of the compound (II). While the reaction can take place in the absence of a solvent, it is generally carried out in the presence of a solvent. The solvent may be, for example an organic solvent capable of dissolving steroid compounds such as an ether (e.g. diethylether, tetrahydrofuran), a halogenated hydrocarbon (e.g. dichloromethane, chloroform, chlorobenzene, dichloroethane, trichloroethylene), an ester (e.g. ethyl acetate, butyl acetate), nitrobenzene, dimethylformamide, dimethylsulfoxide or hexamethylphosphoramide. The reaction is generally conducted within the temperature range of from -10°C to 250°C, when no solvent is employed, or at a temperature within the range of from -10°C to the boiling point of the solvent when a solvent is employed. Following the reaction, the reaction mixture may be immediately treated with water to recover the desired compound. Where R3 is an aralkyl group, the cleavage reaction according to this invention may be carried out by subjecting the compound (II) to catalytic reduction or hydrolysis. The catalytic reduction may be carried out in the presence of a catalyst such as platinum oxide, palladium or Raney nickel, generally in a solvent such as methanol, ethanol, ether or tetrahydrofuran at a temperature within the range of from 10°C to 60°C., and at a pressure within the range of from 1 to 100 kg/cm2. Where R1 is an unsaturated alkyl group, the conditions chosen should be such that the unsaturated bond will not be reduced, e.g. reduction at normal temperature and atmospheric pressure. The hydrolysis is carried out with the same reagent as the ether-cleavage reagent to be employed where R3 is an alkyl or aryl group, or with a halogenoacetic acid such as trifluoroacetic acid, trichloroacetic acid or monochloroacetic acid under the

same conditions as those employed for the ether-cleavage reaction where R² is an

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	alkyl or aryl group (e.g. as to the solvent, reaction temperature and other	
5	Where R ³ is an acyl group, that is where —OR ³ is an esterified hydroxyl group, the cleavage reaction according to this invention may be carried out by subjecting the compound (II) to hydrolysis. This hydrolysis may be conducted by any procedure which enables cleavage of the extendible conducted by any	5
	group in the 3-position without affecting the steroid skeleton or the 16β -alkyl group of the starting compound (II). Thus, for example, the hydrolysis is generally	
10	alcohol (e.g. methanol, ethanol, t-butanol or n-propanol), ether, ethyl acetate, tetrahydrofuran, dimethylsulfoxide or dimethylformamide. The hydrolysis is metal hydroxide (e.g. sodium hydroxide) arganic basic reagent such as an alkali	10
15	potassium carbonate, sodium hydrogen carbonate or potassium hydrogen carbonate), triethylamine or triethylenediamine, or an acid reagent such as an inorganic acid (e.g. hydrochloric acid, sulfuric acid, nitric acid, phosphoric acid) or an organic acid (e.g. formic acid, acetic acid, acetic acid, acetic acid, acetic acid, phosphoric acid) or	15
20	to substantially 80°C.	
20	Where both the R ² and the R ³ groups of the starting compound (II) are acyl groups, both esterified hydroxyl groups in the 3- and 17-positions thereof are generally hydrolysed to free hydroxyl groups, but, if desired, the substituent in the 3-position of the compound (II) may be selectively hydrolysed to convert the esterified hydroxyl group in the 3-position glove to afford the following the 3-position glove to afford the starting compound (II) are acyl groups, but, if desired, the substitution in the 3-position of the compound (II) are acyl groups and 17-positions thereof are 3-position of the compound (II) are acyl groups and 17-positions thereof are 3-position of the compound (II) are acyl groups and 17-positions thereof are 3-position of the compound (II) are acyl groups.	20
25	choosing a mild set of hydrolysing conditions, for example at a comparatively low temperature, e.g. room temperature, using a weakly basic reagent such as an alkali metal carbonate or alkali metal hydrogen carbonate	25
30	rollowing the cleavage reaction of this invention, the contemplated end compound (I) may be isolated and purified by procedures which are conventional per se (e.g. treatment with water, extraction, concentration, recrystallization, chromatography).	30
35	The resulting compounds (I) have an antiestrogen activity, i.e. an inhibitory activity on the binding of estradiol to the estradiol-receptor protein isolated from the tissues of uterine, ovarian or breast carcinomas in mammals including mouse, rat and man, and have substantially no estrogen activity and no androgen activity. Further the present compounds (I) are low in toxicity, and therefore, are of use as antiestrogen drugs for the alleviation of highly estrogen-dependent diseases (e.g. functional uterine haemorrhage, mastitis, breast cancer, uterine cancer) in the said mammalian animals including mouse, rat and cancer.	35
40	Thus for example, the 16\beta-ethylestradiol has an antiestrogen activity which is several times as potent as that of clomiphene and testosterone, and can be used as an antiestrogen drug for the said manufacturing including	40
45	Compounds (I) except for 16β-ethylestradiol may also be employed, depending on the potency of their antiestrogen activity, as antiestrogen drugs in the same manner of usage as testosterone for the alleviation of the above diseases.	45
50	carrier (e.g. lactose, calcium phosphate, corn starch, methyl cellulose, coconut oil, sesame oil, peanut oil) in such dosage forms as tablets, capsules, powders, suspensions or injections.	50
55	These injections may be prepared, for example, by dissolving or suspending the compounds (I) in vegetable oils (e.g. sesame oil, cottonseed oil, castor oil, olive oil, corn oil, peanut oil) in combination, if desired, with antiseptics (e.g. benzyl alcohol, benzyl benzoate, chlorobutanol), solubilizing agents or surface-active agents. Among the compounds (I), 17β-ester derivatives are readily soluble in oils and exhibit a relatively sustained anti-estrogenic action. When the compounds (I)	55

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agents. Among the compounds (1), 1/p-ester derivatives are readily soluble in one and exhibit a relatively sustained anti-estrogenic action. When the compounds (I) are administered orally, they may be administered as powders, tablets, capsules, pills, liquids, syrups, elixirs, buccals or granules. Some example of prescription in which the compounds of this invention are utilized as antiestrogen drugs are given

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For example, where the compound (I) is administered parenterally as an antiestrogen drug for the alleviation of breast cancer, the intramuscular dose range is between 10 and 400 mg, more preferably between 30 and 100 mg, for an adult

			led into 2 to 3 weekly doses of the	
5 .	corresponding smaller amounts. The compound (1) wherein R ² is an acyl group, i.e. 17-ester of 16\beta-alkyl estradiol (I) is, generally speaking, long-active, slow-active, stable in storage and/or easy to prepare in dosage forms in comparison with the 17-hyroxyl compound corresponding thereto. There may be exemplified compositions in which a compound of this invention			5
		ntiestrogen drug;	The a composite of this invention	
	Injectio		10 mainh4 manta	10
10	(1)	sesame oil	10 weight parts 1000 volume parts	10
	(2)		100	
		17-acetate benzyl benzoate	100 weight parts 20 volume parts	
15		sesame oil	1000 volume parts	15
	Capsule			
	·	l6β-ethylestradiol	00 1-14 4-	
		17-acetate lactose	20 weight parts 140 weight parts	
20		corn starch	50 weight parts	20
		sugar ester	4 weight parts	
		calcium salt of	4 maisht sarta	
		carboxymethylcellulose magnesium stearate	4 weight parts 2 weight parts	
25			(220 mg/capsule)	25
	Tablets	•		
	140.00	16β-ethylestradiol		
		17-acetate	20 weight parts	
30		lactose corn starch	100 weight parts 90 weight parts	30
		sugar ester	4 weight parts	
		calcium salt of	A *- \$- \$- A A	
		carboxymethylcellulose magnesium stearate	4 weight parts 2 weight parts	
35		magnostam stout ato		35
33			(220 mg/tablet)	33
	In the pres corresponds to		onds to "gram", and "volume part"	
	The starting	ng compound (II) for this inventi	on may be produced by the method	
40	2100319 0 or h	te specification of German Paters the method described in Che	ent Application As Laid-Open No. mical Pharmaceutical Bulletin Vol.	40
	21, 1393 (1973)), or a method analogous with	the latter method, from the estra-	
	1,3,5(10)-trien-	16-oxo-17β-ols corresponding	to the compound (II) or the	
	generally, the	. cstra-1.3.5(10)-trien-16-oxo-17	2107 (1974). It should be noted that, 3-ols or their derivatives may be	
45	produced by a	procedures similar to the proc	edures established for the species	45
	known among		R2' and R2 are the same acyl group,	
			(I) wherein R ² is hydrogen with an	
50			edures established for the acylation	
50	of the alcohol	lic hydroxyl group. The acyla	ting agent is exemplified by acid onic anhydride, phenylpropionic	50
	anhydride)-org	anic or inorganic bases, acid hal	lides (e.g. acetyl chloride, propionyl	
	chloride, phen	ylpropionyl chloride, benzoyl c	hloride)-organic or inorganic bases,	
55	acias-denyara dicyclohexylca	ting agents such as sulf	uric acid, hydrochloric acid, cylating reaction may be conducted	55
	in the presence	of a catalyst which may be an al	kaline catalyst such as, for example,	,,
	pyridine, picoli	ine, collidine, quinoline or a tert	tiary amine, e.g. triethylamine, or an	
			, e.g. boron trifluoride, zinc chloride or potassium hydrogen sulfate. The	
	or aranamian (oniorido, p-totaque sunonte aciu	or potassium njurogen sunate. The	

(2) To a solution of 0.25 g of 16β -ethylestradiol 3,17-diacetate in 15 ml of methanol is added a solution of 19 mg of anhydrous potassium carbonate in 2 ml of methanol and the mixture is stirred at room temperature for 15 minutes. The

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	reaction mixture is concentrated under reduced pressure and made acidic with 2N-hydrochloric acid, whereupon crystals separate out. Recrystallized from ether-n-hexane (1:1), 16\beta-ethylestradiol 17-acetate is obtained as colourless needles melting at 187 to 188°C.	
5	IR $p_{\text{max}}^{\text{COr}} \text{ cm}^{-1}$: 3400 (OH), 1725 (OCOCH ₃),	5
	Elemental analysis, for C ₂₂ H ₃₀ O ₃ Calcd. C, 77.15; H, 8.83 Found C, 77.19; H, 8.80	
10	Example 5 (1) 16-Ketoestradiol 3-benzylether is reacted with ethyl magnesium iodide in ether to give 16β -hydroxyl- 16α -ethylestradiol 3-benzylether. The product is treated with pyridine-acetic anhydride to give 16β -hydroxyl- 16α -ethylestradiol 3-benzylether 17-acetate. The resulting 17-acetate is heated with zinc powder in toluene at 130° C for 5 hours to give 16β -ethylestrone 3-benzylether. The product is	10
15	treated with sodium borohydride in methanol, whereupon 16β-ethylestradiol 3-benzylether is produced. (2) In 30 ml of methanol is dissolved 0.73 g of 16β-ethylestradiol 3-benzylether,	15
20	followed by the addition of 210 mg of platinum oxide. The catalytic reduction is thus conducted at atmospheric pressure and room temperature. After the absorption of hydrogen has been completed, the platinum oxide is filtered off and the filtrates are concentrated under reduced pressure. By the above procedure, 16β-ethylestradiol is obtained as crude crystals. This crude product is recrystallized from ethyl acetate as in Example 1. In melting point and IR spectrum, this product is in agreement with the product obtained in Example 1.	20
25 30	Example 6 To a solution of 0.93 g of 16β -isopropylestradiol 3-methyl ether in 15 ml of ether is added an ethereal solution of methylmagnesium iodide. The mixture is then treated in the same manner as Example 2, whereupon 16β -isopropylestradiol is obtained as crude crystals. The resulting crude crystals are recrystallized from ethyl acetate. Melting point: 221 to 222°C.	25 30
	IR $v_{\text{max}}^{\text{Kbr}}$ cm ⁻¹ : 3400 (OH), 1610, 1590 (Ar).	
	NMR $\delta_{ppm}^{d_0-OMSO}$: 0.70 (3H, s, 18-CH ₃), 0.83 (3H, d, J=5 Hz, CH ₃), 0.98 (3H, d, J=5 Hz, CH ₃), 3.73 (1H, d, J=9 Hz, 17 α -H), 6.4—7.2 (3H, m, Ar)	
35	Elemental analysis, for $C_{21}H_{20}O_{2}$ Calcd. C, 80.21; H, 9.62 Found C, 80.30; H, 9.67	35
40	Example 7 Under ice-cooling 0.2 g of phosphorus tribromide is added in small portions to a solution of 0.6 g of 16β -ethylestradiol 3-methyl ether in 10 ml of dichloromethane. The resulting mixture is allowed to stand at room temperature for 4 hours. The reaction mixture is poured in small portions into ice-water and extracted with	40
45	dichloromethane. Upon removal of the solvent by concentration, 16β -ethylestradiol is obtained as crude crystals. Recrystallization under the same conditions as in Example 1 yields pure crystals. In melting point and IR spectrum, this product is in agreement with the product obtained in Example 1. In a similar manner to the above, 16β -allylestradiol is obtained from 16β -allylestradiol 3-methyl ether. Melting point: 204 to 206°C.	45
	IR $\nu_{\text{max}}^{\text{KBr}}$ cm ⁻¹ : 3350 (OH), 3080, 1640 (allyl), 1610, 1595 (Ar).	
50	Elemental analysis, for C ₂₁ H ₂₆ O ₂ Calcd. C, 80.73; H, 9.03 Found C, 80.77; H, 9.10	50
	Example 8 (1) To a solution of 0.3 g of 16β-ethylestradiol in 2 ml of pyridine is added 0.6	55

5	ml of propionic anhydride. After keeping the resulting mixture at 50° C for 10 hours, 10 ml of water are added to the reaction mixture, followed by extraction with dichloromethane. The organic layer is washed with water, dried over anhydrous sodium sulfate and concentrated, whereupon crude crystals are obtained. Recrystallization from methanol gives 16β -ethylestradiol 3,17-dipropionate as colourless needles melting at 57° C.	. 5
	IR $p_{\text{max}}^{\text{KBr}} \text{ cm}^{-1}$: 1760 (OCOC ₂ H ₆), 1725 (OCOC ₂ H ₆).	
10	(2) To a solution of 0.2 g of 16β -ethylestradiol 3,17-dipropionate in 10 ml of methanol are added 16 mg of anhydrous potassium carbonate, followed by stirring at room temperature for 30 minutes. The reaction mixture is concentrated under reduced pressure, and the residue is made acidic with 2N-hydrochloric acid, whereupon crystals are obtained. The crystals are collected by filtration and recrystallized from hexane to give 16β -ethylestradiol 17-propionate as colourless needles melting at 176 to 178 °C.	10
15	IR $\nu_{\text{max}}^{\text{KBr}} \text{ cm}^{-1}$: 3350 (OH), 1700 (OCOC ₂ H ₈).	15
	Elemental analysis for C ₂₅ H ₃₂ O ₃ Calcd. C, 77.49; H, 9.05 Found C, 77.48; H, 9.07	
20	Example 9 (1) In a similar manner to Example 4-(1), 16β-isopropylestradiol 3,17-diacetate is obtained by acetylation of 16β-isopropylestradiol with acetic anhydride-pyridine. Melting point: 115 to 116°C.	20
	IR $p_{\text{max}}^{\text{KBr}} \text{ cm}^{-1}$: 1765 (OCOCH ₂), 1735 (OCOCH ₂).	
2 5	(2) In a similar manner to Example 4-(2), 16β -isopropylestradiol 3,17-diacetate is hydrolysed with anhydrous potassium carbonate to give 16β -isopropylestradiol 17-acetate. Melting point: 193 to 194°C.	25
	IR $\nu_{\text{max}}^{\text{KBr}} \text{ cm}^{-1}$: 1350 (OH), 1700 (OCOCH ₃).	
30	Elemental analysis for $C_{23}H_{22}O_3$ Calc. C. 77.49; H, 9.05 Found C, 77.31; H, 9.11	30
35	Example 10 (1) To a solution of 0.2 g of 16β-ethylestradiol 3-methyl ether 17-acetate in 10 ml of dimethylsulfoxide is added 0.5 g of dried sodium iodide, and the mixture is refluxed for 3 hours under a nitrogen gas stream. After cooling, 30 ml of water are added to the reaction mixture, and the resulting mixture is extracted with ether. The ether layer is washed with water, dried over anhydrous sodium sulfate and concentrated, whereupon pale yellow crude crystals are obtained. Recrystalization from ether-hexane (1:1) gives 16β-ethylestradiol 17-acetate. This product is in accordance with the product obtained in Example 4 in melting point	35
40	and IR spectrum. (2) Similarly to Example 7, 16β-ethylestradiol 3-methylether 17-acetate is treated with phosphorus tribromide to yield 16β-ethylestradiol 17-acetate.	40
45	Example 11 (1) To a solution of 0.3 g of 16β -ethylestradiol in 10 ml of pyridine is added 0.5 g of 3-phenylpropionyl chloride, and the mixture is kept at room temperature for 12 hours. 10 ml of ice-water are added to the reaction mixture and the mixture is extracted with ether. The ether layer is washed with a 3N-aqueous solution of potassium carbonate, dried over anhydrous sodium sulfate and concentrated, whereupon 16β -ethylestradiol 3,17-diphenylpropionate is obtained.	45
50	IR $\nu_{\text{max}}^{\text{Neat}}$ cm ⁻¹ : 1760, 1735 (OCOCH ₂ CH ₂ —C ₆ H ₅).	50
	(2) To a solution of the product obtained in the above experiment (1) in 10 ml of methanol is added 0.1 g of potassium carbonate and the mixture is stirred at	

(2) In a similar manner to Example 2, 16β -n-butylestradiol 3-methylether is reacted with methylmagnesium iodide to give 16β-n-butylestradiol melting at 148 to 150°C (recrystallization from hexane).

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IR $v_{\text{max}}^{\text{KBr}} \text{ cm}^{-1}$: 3400 (OH), 1605 (Ar).

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Elemental analysis for C₂₂H₃₂O₂
Calcd. C, 80.44; H, 9.83
Found C, 80.40; H, 9.99 45

Mass: m/e 340 (M⁺), 325 (-15), 322 (-18).

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In a similar manner to the above experiment (2), 16β -(3-butenyl)-estradiol is obtained from 16β-(3-butenyl)estradiol 3-methylether.

Melting point: 154 to 156°C.

IR $\nu_{\text{max}}^{\text{KBr}}$ cm⁻¹: 3400 (OH), 3050, 1635 (c=c), 1605 (Ar).

Elemental analysis for C₂₂H₃₀O₂
Calcd.
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WHAT WE CLAIM IS:— I. A compound of the formula (I):

wherein R1 is an alkyl group or an alkenyl group of two or more carbon atoms, and

wherein R' is an alkyl group or an alkenyl group or two or more carbon atoms, and R² is hydrogen or an acyl group (as herein defined).

2. A compound as claimed in Claim 1, wherein the alkyl group represented by R' is a lower alkyl group having 2 to 4 carbon atoms.

3. A compound as claimed in Claim 1 or 2, wherein R² is hydrogen.

4. A compound as claimed in Claim 1 or 2, wherein R² is an acyl group.

5. A compound as claimed in Claim 4, wherein the acyl group represented by P² is lower alkylearbonyl whose alkyl mojety is alkyl having 1 to 3 carbon atoms 15

R² is lower alkylcarbonyl whose alkyl moiety is alkyl having 1 to 3 carbon atoms, benzoyl or phenylpropionyl.

6. 16β-ethylestradiol.
7. 16β-ethylestradiol 17-acetate. 20

16β-isopropylestradiol.
 16β-allylestradiol.

10. 16β-ethylestradiol 17-propionate. 25

11. 16β-isopropylestradiol 17-acetate.
 12. 16β-ethylestradiol 17-phenylpropionate.
 13. 16β-ethylestradiol 17-benzoate.

14. 16β-n-butylestradiol.

15. 16β-(3-butenyl)-estradiol.
16. A pharmaceutical composition comprising any one of the compounds claimed in Claims 1 to 15, together with a pharmaceutically acceptable carrier or diluent therefor.

17. A process for producing a compound of the formula (I)

wherein R' is an alkyl group or an alkenyl group of two or more carbon atoms, and 35 R² is hydrogen or an acyl group (as herein defined), which process comprises subjecting a compound of the formula (II):

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wherein R¹ has the same meaning as defined above, R^{2'} is hydrogen or an acyl group (as herein defined and R³ is a hydrocarbon radical or an acyl group (as herein

defined), to cleavage of the acyl group or hydrocarbon radical or an acyl group (as herein defined), to cleavage of the acyl group or hydrocarbon radical of the etherified or esterified hydroxyl group in the 3-position thereof.

18. A process as claimed in Claim 17, wherein R³ is an acyl group.

19. A process as claimed in Claim 17, wherein R³ is a hydrocarbon radical.

20. A process as claimed in Claim 19, wherein the hydrocarbon radical represented by R³ is lower alkyl having 1 to 3 carbon atoms, phenyl, p-nitrophenyl, beautyl or hearthydryl. benzyl or benzhydryl.

21. A process as claimed in Claim 18, wherein the acyl group represented by R3 is lower alkylcarbonyl whose alkyl moiety is alkyl having 1 to 3 carbon atoms, or arylcarbonyl.

22. A process for producing a compound (I) as defined in Claim 1, substantially

as herein described with reference to any of the specific examples.

23. Compound (I) as defined in Claim 1 when produced by a process as claimed in any of Claims 17 to 22.

24. A pharmaceutical composition comprising at least one compound (I) as claimed in Claim 23, together with a pharmaceutically acceptable carrier or diluent therefor.

> **ELKINGTON & FIFE,** Chartered Patent Agents, 52-54 High Holborn, High Holborn House, London WC1V 6SH. Agents for the Applicants.

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